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- Applicant: CHEMEX PHARMACEUTICALS, Inc. 1401-17th Street, Suite 850
 Denver, Colorado 80202(US)
- ② Inventor: Allen, Larry M. 450A Josephine Street Denver, Colorado 80206(US)
- Representative: Allard, Susan Joyce et al BOULT, WADE & TENNANT, 27 Furnival Street London EC4A 1PQ(GB)

(A) Pharmaceutical vehicles for reducing transdermal flux.

This invention relates to the use of a water-soluble part of compound in a topical pharmacoutical compound in a topical pharmacoutical composition containing a pharmacologically active agent to enhance the skin or mucous pentarian penetration and retention of the pharmacologically active agent, and to reduce the transfermal flux through the skin and mucous membrane and retention of the pharmacologically active agent, and to reduce the transfermal flux through the skin and mucous membrane.

Field of the Invention

Method of inducing reservoir effect in skin and mucous membranes so as to increase penetration and residence time of pharmacologically active agents therein.

Background of the Invention

There are many localized disease conditions which are effectively treated by topical application of suitable physiological agents. In order for such treatments to be maximally effective, it is necessary that as 10 much of the pharmacologically active agent as possible be absorbed into the skin where it can make contact with the disease condition in the dermal fissues without being lost by rubbing off on clothing or evaporation. At the same time, the agent must not penetrate so effectively through the skin as to be rapidly lost to the lymphatic and vascular circulatory systems. This latter factor is especially important when the pharmacologically active agent is toxic when used systemically.

The ideal vehicle for topically applied pharmaceuticals is therefore one which can produce a "reservoir effect" in the skin or mucous membranes to which the topical treatment is applied. This "reservoir effect" is defined as an enhancement of the skin or membrane's ability to both absorb and retain pharmacologically active agents, i.e., to increase skin or membrane residence time, decrease drug transit time and reduce transfermal flux.

20 A number of compounds are known to enhance the ability of pharmacologically active agents to penetrate the skin and mucous membranes, for example, N-bis-azacyclopentan-2-ony-laikones, 1-substituted azacyclobeptan-2-ones and higher alkyl-substituted azacyclopentan-2-ones, as well as directivisation ide and lower alkyl sufloxides. These compounds, however, have the disadvantage of allowing rapid systemic dispersion of the pharmacologically active agents away from the localized site of pathology, as the part of particular to the control of a proposal medicaments, such as the retinoids used in the treatment of acen, and methorexate, used in the treatment of pendicaments, such as the retinoids used in the retainment of acen, and methorexate, used in the treatment of pendicaments. Thus, there is a reset for a method of enhancing the ability of such medicaments to penetrate into the skin or mucous membrane so that a lesser total dosage may be used, while at the same time retarding their ability to move from the skin to the interior of the body.

The problem posed by the paradoxical requirement that a systemically toxic topical medicament be efficiently absorbed into, but not through, the skin, has heretofore gone unrecognized. There have been no vehicle additives available to the pharmacological industry which act to enhance the "reservoir capacity" of the skin and mucous membranes so that the amount of pharmacologically active agents reaching and being retained at the size of localized pethologies is maximized.

Summary of the Invention

This invention is the use of a water-soluble zinc-containing compound in a topical pharmaceutical composition containing a pharmacologically active agent to enhance the skin or mucous membrane penetration and retention of the pharmacologically active agent, and to reduce the transfermal flux through the skin and mucous membrane.

The additives of this invention are water-soluble zinc-containing compounds, preferably zinc halide, zinc sulfate, zinc nitrate, zinc acetate, and/or zinc stearate, and most preferably zinc chloride.

The pharmacologically active agents with which the water-soluble zinc-containing compounds are used are preferably those containing hydroxyl, oxo, sulfhydryl, amine, carboxyl, and other anionic groups in configurations which readily allow complexation or chelation with zinc ions.

In inducing a reservoir effect in the skin and mucous membranes to reduce transdemal flux so that drugs are absorbed and retained therein in larger amounts for longer periods of time than has herefolore been possible, the water-soluble zinc-containing compounds of this invention act as potentiators for the so pharmacologically active agents. Potentiation is defined as overcoming or reducing undestrable effects such as systemic toxicity and extending the ranse of effectiveness of the pharmacologically active agent, or both,

Description of the Drawing

The drawing is a graph of test results showing the skin reservoir effect achieved by the method of this invention utilizing nordihydrogualeric acid with zinc chloride, as compared to the same compound without zinc chloride. The nordihydrogualeric acid was labelled with carbon-14 for radiotracer analysis. The graph shows immediate absorption and longer restinon of larger amounts of the nordihydrogualeric acid with

zinc chloride present than without. The availability of nordihydroguaiaretic acid, a lipoxygenase inhibitor, to therapeutically act upon localized pathologies, both with and without added zinc chloride, is measured by the areas under the respective curves. It is apparent that stin bioavailability is greatly enhanced in the presence of a zinc-containing compound. The drug flux rate can be calculated as a function of the area of under the curve by dividing this area by dose. As is apparent, drug flux rate is substantially decreased over the entire dosage range by the addition of zinc chloride.

Detailed Description of the Invention

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The zinc-containing compounds of this invention are generally any water-soluble organic or inorganic zinc satts which dissociate in the topical vehicle so as to provide zinc ions for complexation or chelation with the pharmacologically agents present in the vehicle. Examples of suitable zinc-containing compounds are zinc halide, zinc sulfate, zinc nitrate, zinc acetate, and/or zinc stearate. The most preferred zinc-containing compound is zinc chloride.

Such water-soluble zinc-containing compounds may be prepared by means known to the art, and many are commercially available.

The topical preparations of this invention having enhanced reservoir inducing capacities may be prepared by mixing a water-solube zinc-containing compound into the pharmacoutical preparation or vehicle in which a reservoir-inducing capacity is to be created by means known to the art so as to promote accomplexation or chelation is the pharmacologically active agents therein with zinc ions. The presence of other metallic ions which would unfavorably complete with zinc for complexation or chelation sites is to be avoided. The zinc salt can also be mixed with the pharmacologically active agent to form a chelate or complex which is then incorporated into the pharmacoutical carrier.

The zinc-containing compounds are preferably present in an equimotar ratio with the pharmacologically as active agents, so as to cause maximum complexation or chelation. Where stratum commeum destruction, i.e. decormification, is desirable, an excess of such zinc-containing compounds which also act as escharotics, e.g. zinc chloride, may be used. (Generally, concentrations of 35% (0.257 moles per 100 grams) or more zinc chloride will cause tissue destruction when topically applied.) Normally, use of equimotal concentrations of zinc chloride and the pharmacologically active agent will not involve the use of escharotic amounts of zinc chloride, however less than 35% zinc chloride should considered an upper limit when no escharotic effect is desired. Less than an equimotar ratio of zinc-containing compound to pharmacologically active agent may be employed where it is not desired that all the medicament be absorbed into the skin or mucous membranes, e.g., in connection with mouthwash and douche preparations where attack on free-swimming organisms is also desirable.

Other ingredients may be added to the preparations, including coloring agents, stability-enhancing agents, antioxidants, and the like. Preferably these additives will not compete with the pharmacologically active agents for zinc; however when necessary, excess zinc-containing compounds may be used to compensate for the zinc complexing or chelating effect of such additives.

The pharmacologically active agents of this invention are those intended for topical application to a chieve localized therapoutic or cosmetic effects. A partial list of suitable pharmacologically active agents includes steroids, antifungals, anti-uricellular microorganism agents, antiviral agents, antiparasitic agents, antinepassitic agents, anti-epostastic agents, anti-epostastic agents, anti-epostastic agents, anti-entral agents, anti-inflammatory agents, immun-opharmacological agents, allergens, antilistaminic agents, anti-inflammatory agents, anneum-opharmacological agents, antiepostastic agents, radiopaque agents, cryoprotective agents, perfures, sinscriptionalisms, bair dyses, antiscarring agents, sun screens, melanin-stimularing agents, antiprespirants, antissecretory agents, depilatories, hair restorers, winkle-reducing agents, antidandruff agents, emollients, rubfacients, and cosmetic agents in general.

The mechanism by which the reservoi-inducing effect of this invention is produced is not known, however, it is preferred that the pharmacologically agents contain hydroxy, oxe, sulfflydry, amino, carboxyl, so or other anionic groups, or combinations thereof, in conformations which allow complexation and/or chelation by sinc ions.

Preferred pharmacologically active agents of this invention are:

Antineoplastic agents including NIDGA (nordihydrogualaretic acid), VP-16 (epipodophyllotoxin beta-otherhyldene glucopyranoside-etopoide), VM-26 (epipodophyllotoxin beta-Otherhyldene glucopyranoside-teriposide), M-26 (epipodophyllotoxin, beta-Otherhyldene glucopyranoside-teriposide), 4'demethyl epipodophyllotoxin, diethylstillostrol, dithranol, cyclophosphamide, millomycia, daunomycin, Dalfrum dis-dishinel-dishindira, dariamycia, lounds-Flusorouracii, and methotrexia.

Immunopharmacological agents which may be topically applied including polypeptide nanoparticles comprising interleuken or active fragments thereof, antibodies or active fragments thereof, interferons, and liposomes. Such delivery systems providing sustained release of pharmacologically active agents are effectively localized or held in place by zinc according to this invention.

- 3. Steroids, which are utilized for a wide range of therapeutic purposes including anti-inflammation, antipuritie, enhancement of moisture retention, etc., including dexamethasone, hydrocordisone sectate, hydrocordisone sectate, hydrocordisone sectate, braincinolore acetoride, purpositione, hydrocordisone exteroide, discondinied, fluorisonione sectoride, discondinied, fluorisonione sectoride, discondinied, fluorisonione discondinied, fluorisonione, discondinied, fluorisonione, and fluorisonione, discondinied, fluorisonione, discondinied, fluorisonione, discondinied, fluorisonione, and fluorisoni
- 4. Antifungal agents which are used to treat fungus infections on the skin, hair, and nails, such as athlete's foot (tinea pedis), lock itch (tinea cruris), ringworm (tinea corporis), which can be caused by a number of fundi, particularly Tricophyten rubrum, Trichophyten mentagrophytes, and Epidermophyten floccosum, and Microsporum canis. These antifungal agents include haloprogin, iodochloro, miconazole nitrate, tolnaftate, thiabendazole, chloroxine, amphotericin, candicin, fungimycin, nystatin, chlordantoin, clotrimazole, ethonam nitrate, miconazole nitrate, pyrrolnitrin, fezatione, ticlatone, tolnaftate, triacetin, carbonic acid derivatives; dithiocarbamate, thiourea, thiocyantes; aromatic carboxylic acids and the amides thereof, benzoic acid, salicylic acid, salicylic acid amide and anilide; aromatic sulfides, polysulfides, and sulfoxides, 5,5-dichloro-2,2-dihydroxydiphenylsulfide; invert soaps, quaternary ammonia and phosphonium compounds, decamethylene-bis-(4-thio-pyridine-methyl-tosylate; quinoline derivatives, 8hydroxyquinoline sulfate, halogenated quinolines, 7-iodo-8-hydroxy-quinoline-5-sulfonic acid, 5-chloro-7iodo-8-hydroxy-quinoline, 5-chloro-8-hydroxy-quinoline, 5,7-dichloro-8-hydroxyquinaldine, 5,7-diiodo-8hydroxyquinoline, decamethylene-bis (4-amino-quinaldium chloride); benzothiazole derivates, (2dimethylamino-6-(beta-diaminoethoxy)-benzothlazole dihydrochloride; imidazole derivatives, 1-(o-chloroalpha-alpha-diphenyl-benzyl)-imidazole, 1-[o,p-dichloro-beta-(o,p-dichlorobenzyloxy)-

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phenethyliumidazole); benzimidazole derivatives, 2-phenylbenzimidazole, 2-furfurylbenzimidazole, indiadizine derivatives, 35-dibenzipidatyndy-indiadizine derivatives, 3-furfuz-dipteratynd-indiadizine derivone, tran derivatives, 5-furfuz-diviny-3-chloropropionate; quinones, tetrachloro-p-benzoquinone, 1.4-naphthoquinone, phenarthraquinone; sulnomarides and suffores zonamida diadinidines, 2-furfuzovstilibandine, and diamitinical ordinamidation derivatives and diamitinical diadinidines.

Antifungal agents are also used to treat vaginal infections caused by Candida albicans and related yeasts; these gents include directly sodium sulfusuccinate, hadprogin, microargice intritiee, potassium sorbate, propionate compounds, such as calcium propionate and sodium propionate, and sodium lauryl sulfate.

- 5. Antibacterial agents, which are utilized for treating skin infections such as impetigo, ecitymus, follicultitis, boils, and acute pronychia, and for treating skin wounds and as a wound cleanser, and which may be used in this invention, including sulfonomides, penicililins, cephalosporins, penicililinse, lincomycins, vancomycins, letracylines, chloramphenicols, and streptomycins; including within this group the following compounds: grandictin, neomycin, polymynis beta sulfate, letracycline, beneathonium chloride, gentamicin sulfate, nitrofurazone, benzalkonium chloride, gentamicin sulfate, nitrofurazone, benzalkonium chloride, pentamicin sulfate, nitrofurazone, pentamicin sulfate, nitrofurazone,
- 6. Antiviral agents, including those used to treat warts, such as glacial acelic acid, ascorbic acid, calcium partotherate, lactic acid, salicipic acid, canthardin, and podophylin; and antiviral agents used to treat cold sores or herpes simplex such as ecyclovir, benzalikonium chloride, alcohol, allarotio, anytydrous glycerin, benzocaine, camphor, carbamide peroxide, fanolin, memthol, pertodum, and phenot, and antiviral agents including those used to treat herpes genitalis such as urea, idoxuridine, amantadine, methisazone, cvirarabine, indreferors, chloroform, ether, bacillus calmete-querin, and levamisolo.
- Antiparasitic agents including antihelmintic agents (agents that destroy or expel intestinal worms) capable of penetrating the skin of the animal to be treated, e.g. benzimidazole compounds, tetramisole, levamisole, and iscoulinoline compounds.
- Pediculicides, for mites (or scables) and lice, including lidane, pyrethrins, piperonyl butoxide, malathion, and crotamiton.
- 9. Acne treatment compounds inclusting benzoyl perovide, resorcinol, resorcinol monoacetats, sulfur, povidone-rioline, salicytic acid, phenol, fluoriolone acetenide, para-aminobenzoi acids, sodium thiosulfate, meciocyline sulfosalicylate, sodium sulfacetamide, tetracycline hydrochloride, aliphatic dicarboxylic acids, acid, acid, and sulfurated lime.
- 10. Antipsoriasis agents including cytostatic agents, which retard skin-cell growth, keraldytic agents, which loses and dissolve scales, tar preparations, whose mode of action is uncertain; hydrocordisone preparations, which reduce litching and inflammation; anti-tich preparations; and antimicrobials. These antipsoriasis agents include coal tar preparations, jumiper tar, pine tar, allantion, sapponated cresol, menthol, mercury oleate, phend preparations, resortion, salicytic acid, antimatin, and methotrexate.

- 11. Leprosy agents including 4-4'-diaminodiphenyl sulfone.
- 12. Anesthetic agents for pain and itching, Inflamed skin, sunburn, insect bites, burns, wounds, hemorrholds, poison ivy, poison oak, including: benzocaine, lidocaine, lidocaine, lydrochloride, dibucaine, dydrochloride, procaine, tetracaine, tetracaine, betracaine hydrochloride, promotine, dyclonine, hydrochloride, pramoxine hydrochloride benzyl alcohol, diperodon, butamben picrate, cyclomethycaine sulfate, and dimentilisouin hydrochloride.
- 13. Analgesic agents for pain and litching, inflamed skin, sunburn, insect bites, burns, wounds, hemorrholds, poison ivy, poison oak, including: salicylic acid derivatives; N,N-dimethyl aspartic acid; N-N-dimethyl alutamic acid; trolamine salicylate, methyl salicylate; antipyrine, asprini, and salicylamide.
- 14. Counter-irritants (agents applied locally to produce an inflammatory reaction with the object of listracting and relieving a doep seated inflammatory process) including methyl salicylate, campor, menthol, eugenol, eucalyptol, thymol, slayl slothicocyanate (mustard oil), capsicum preparations, histamine dithordorchloride, methyl inclorate, and turpentine oil.
- Antihistamines which are used principally against itching, but are also mildly anesthetic, including diphenhydramine hydrochloride, phenyltoloxamine dihydrogen citrate, pyrilamine mleate, tripelennamine hydrochloride.
 - 16. Diagnostic agents including altergenic extracts for diagnosis and immunotherapy of specific altergy offenders from the following categories: pollens, foods, dusts, epidermals, insects and stinging insects, fungl, molds, yeasts; tests for sensitivity to therapeutic periclifilin (benzyl-poincillipy)-polysting); tests for sensitivity to tetanus antigens, diphtheria antigens, streptococcus antigens, tuberculin, Candida antigens, Trichophyton antigens, and Proteous antigens.
 - 17. Vitamins and nutrients for skin, hair, and scalp conditions, including anti-scarring agents, vitamins B_8 , B_6 ,
 - 18. Cosmetic agents and perfumes including compositions to reduce the appearance of wrinkles such as water soluble elastin and pregnenolone; skin depigmenting agents and bleaches, including hydroquinone and monobenzone.
 - 19. Sunscreens including: dioxybenzone, oxybenzone, padimate O, padimate A, aminobenzoic acid, cinoxate, diethanolamine p-methoxycinnamata, et why 4-tbistyncyopropyll aminobenzoate, ethylhexyl salicytate, glycenyl aminobenzoate, shymbol common and dithydroxyacetone, red petrolatura, and suissohenzone.
 - The foregoing are simply examples of pharmacologically active agents including therapeutic and cosmetic agents which may be used with enhanced effectiveness for their known properties in accordance with this invention
- In addition to the reservoir effect produced by zinc ions in skin and mucous membranes, a direct potentiating effect has been observed when pharmaceutical preparations containing zinc ions and pharmacologically active agents are injected directly into diseased tissues, particularly solid tumors. The mechanism for this potentiating effect is not known; however, it may be caused by a reservoir-inducing effect directly on the tissues involved.
- Dosage forms for topical application may include lotions, cintments, creams, gels, suppositories, nasal solutions, mouthwashes, sprays, acrosols and the like. Typical carriers which make up the foregoing dosage forms include water, acetone, isopropyl alcohol, stearyl alcohol, froms, ethyl alcohol, polyvniy pyrmolione, propylene glycol, polyethlyene glycol, fragrances, gel-producing materials, mineral oil, stearic acid, spermaceti, sorbitan, monoleate, polysorbates, "Tweens", sorbitan, fragrances?
- The amount of the composition, and thus of the pharmacologically active agent therein to be administrated, will be ascertained by the practitioner utilizing his ordinary skill. Due to enhanced activity which is achieved, the dosage of agent may often be decreased from that generally applicable. In accordance with usual prudent formulating practices, a dosage near the lower useful range of the particular agent may be employed intallity and the dosage increased as indicated from the observed response.

Preferred compositions are illustrated in the following examples:

EXAMPLE 1

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Antineoplastic Preparation

Two compositions utilizing zinc chloride, nordihydroguaiaretic acid (NDGA), acid (EDTA), butylated hypotylolulene (BHT), stearyl alcohol, purified water, polyethylene gylcol having an average molecular weight of 400 (PEGO 400), and polyethylene glycol having an average molecular weight of 3350 (PEGO).

3350) were prepared in the following manner: the purified water was placed in a clean glass container of suitable capacity; the water was heated to about 80-90°C with stirring; and zinc chloride was added to the heated water, continuing the stirring until the zinc chloride dissolved. The ethylenediaminetetracetic acid was next slowly added with mixing until dissolved. In a separate glass container of suitable size, the polyethylene glycol 400 was heated to about 80-90°C with stirring; the NDGA was added thereto; then the BHT; and this mixture was then cooled to about zone temperature and passed through a number 3 cilloler mill until smooth. The polyethylene glycol 3350 was then heated to about 80-90°C in a suitable container and the milled incredients added thereto with mixture.

The final compositions in wt/wt % were as follows:

TABLE 1

Composition	Com	pound
	Α	В
zinc chloride	29.8	10.0
NDGA	4.6	4.6
EDTA	14.7	4.93
BHT	1.1	1.1
stearyl alcohol	0.5	0.5
H ₂ O	18.3	18.3
PEGO 400	26.4	26.4
PEGO 3350	4.5	4.5

EXAMPLE 2

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30 Reservoir Effect

This study was designed to provide basic pharmacokinetic date on the disposition of Carbon 14 (¹⁴ C) label on ordiflydroguainetic scif (NDGA) applied dermally in modified compounds A and B described in Example 1. In addition, the distribution of zinc was measured for the dermally applied Compound A.

The ¹⁴C-NDGA compound exhibited a specific radioactivity of 20.2 Ci/mole (66.9 micro Ci/mg) and a purity of 96.9% by mass spectrometry and by radioautography of thin-layer chromatography plates developed in benzeneisopropanotacetic acid-water (255:210).

Subsequently, 25.1 mg of the ¹⁴C-NDGA-Compound (66.9 micro Ci/mg) were mixed with 12.35 g of Compound A. Analyses of triplicate samples of the final mixture for ¹⁴C by counting and for NDGA by high-pressure liquid chromategraphy (PHC) demonstrated homogeneity, with a content of 51 micro g of ¹⁴C-NDGA compound/mg of Compound A. The specific radioactivity of the NDGA was 3.00 x 10³ micro Ci/micro

Similarly, 26.3 mg of the original "C-NDGA compound (66.6 micro Climg) were mixed with 12.55 g of Compound A devoid of Z and EDTA to obtain a mixture for the study of the dermal penetration of NDGA from Compound A devoid of Z and EA. Analyses of triplicate samples of the modified Compound A showed the final mixture to be homogeneous with regard to "C and NDGA; it contained 55 micro g of "C-NDGA compounding of whelice. The specific radioactivity of the distude NDGA was 3.41 x 10⁻³ micro Climicro g.

The compounds were dermally applied to young adult Syraque-Dawley rats by the following protocol: under ether anesthesia, the back skin of the rat was prepared by removing the hair from a 5 x 5-cm area so with a clipper and the residual hair stubble was removed with a wax depliatory. Then the skin was stripped repeatedly (5x) with adhesive tape until the statum comeum was removed. Then 0.5 gm of the formulation was weighted on a 5 x 5-cm sheet of polypropylene, which was applied to the prepared skin. It was secured in place by hypoallegenic lape. Finally, the bandage was overwapped with bandage taps. Alter treatment, the rats were caped individually in metabolism cages, which allowed free access to food and water and provided for separate collection of urine and faces.

The testing of Compound A with ¹⁴C-NDGA was performed in 15 male Sprague-Dawley rats (mean weight 339 ± 16 g). They received an average of 0.520 (± 0.032) g of Compound A containing ¹⁴C-NDGA. The mean dose of ¹⁴C-NDGA was 78.5 (± 7.0) mg/kg of body weight. The rats were housed individually in

metabolism cages providing for free access to food and water and for the separate collection of urine and feces. Groups of 3 rats were sacrificed at 4, 24, 48, 72, and 96 hr and excreta were collected from each rat during 24-hr periods. In addition to the usual collection of tissues, the skin site of application was excised after wiping the site with water-moistened tissue. The wipes were added to the wrappings, which were 5 immersed in a small container of acetone.

The testing of Compound A devoid of Zn and EDTA was performed on 15 male Sprague-Dawley rats (mean weight 241 ± 7 g). They received an average of 0.390 (± 0.019) g of Zn-free C205 containing 14 C-NDGA. The average dose of 14 C-NDGA was 83.2 mg/kg of body weight. Groups of three rats were bled terminally and tissues were taken at 4, 24, 48, 72, and 96 hr after dosing. At each sacrifice time, those three 10 rats scheduled to be sacrificed next were also bled nonterminally from the orbital sinus. The wrappings and wipes of the skin site were taken at the time of sacrifice and added to acetone as described above.

The results of the study are given in Table 2A for Compound A containing 14 C-NDGA and Table 2B for Zn-Free Compound A containing 14 C-NDGA. The results of analysis for tissue distribution of zinc as a percent of dose in rats receiving Compound A containing 14 C-NDGA are given in Table 2C.

TABLE 2A

	Tissue Distribution of ¹⁴ C as a Percent of Dose in Rats Receiving Compound A containing ¹⁴ C-NDGA, Dermally								
Ī	Tissue		se of 14 C Found at I	C Found at Hours					
		4	24	48	72	96			
	Organs (%) Skinsite (%)	4.62 (±2.27) 20.8 (±8.4)	7.55 (±1.85) 19.6 (±5.2)	10.29 (±8.8) 11.1 (±1.4)	10.43 (±6.94) 8.58 (±5.41)	13.45 (±3.45) 7.01 (±2.28)			

 $^{9}N = 3$

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TABLE 2B

	Tissue Distribution of ¹⁴ C as a Percent of Dose in Rats Receiving Zn-Free Compound A Containing ¹⁴ C-NDGA, Dermally								
Γ	Tissue	,	Mean (± S.D.) ^a Percent of the Dose of ¹⁴ C Found at Hours						
		4	24	48	72	96			
Γ	Organs (%)			10.47 (±10.33)	5.41 (±4.21)	8.03 (±1.48)			
L	Skinsite (%)	4.15 (±0.76)	12.20 (±5.9)	7.86 (±3.75)	4.72 (±3.18)	1.43 (±0.55)			

TABLE 2C

Tissue Distributio	n of Zn as a Per	cent of Dose	in Rats Receiv	ing Compound	A, Dermall		
Tissue	Mean Percent of the Dose of Zn found at Hours						
	4	24	48	72	96		
Organs (%) Skinsite (%)	3.28 10.5	6.54 11.8	6.79 9.58	11.47 6.46	12.69 4.99		

 $^{9}N = 3$

The study was continued for testing Compound B with 16 C-NDGA; Zn- and EDTA-free Compound B; and modified Compound B with no BHT and 0.10 EDTA. The following Table 2D lists the compositions of the compounds and the amounts of materials used for preparing the compounds containing 14C-NDGA.

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These compounds were analyzed for ¹⁴C by scintillation counting and for NDGA by liquid chromatography.

TABLE 2D

5			Composition (1) Modified
	Constituent	Compound B Formulation	Compound B Formulation	Compound B Formulation
10	Compound Zn Compound EDTA Compound NDGA Compound BHT	10.00 4.93 4.60 1.10 18.32	0.00 0.00 4.60 1.10 18.32	10.00 0.10 4.60 0.00
15	Water, purified PEG 400 PEG 8000 Stearyl Alcohol	2.60 53.45 5.00	14.50 49.39 12.09	19.42 14.19 46.69 5.00
20	Compounds containing	ng 14C-NDGA:		
	mg of 14C-NDGA	25.75	25.40	25.20
25	g of Formulation	12.55	12.55	12.75
	NDGA in final mixture	4.80	4.79	4.78

The mean rat body weights, average doses of the formulations, and mean doses of "C-NDGA in mg/kg of body weight for the three current protocols were: 297 ± 15 (standard deviation), 512 ± 28 mg, and 82.7 ± 2.0 mg/kg for Compound B; 325 ± 12 g, 570 ± 28 mg, and 84.0 ± 1.4 mg/kg for Zn-free Compound B; and 328 ± 27 g, 575 ± 45 mg, and 84.2 ± 28 mg/kg for modified Compound B. Filteen rats were used for each study and groups of three rats were sacrificed at 4, 24, 48, 47, 22 and 89 f. artien, 48, 72 and 89 f. artien, sees, and compound says ings were collected. Also, from the groups sacrified at 24, 48, 72, and 89 fr. urine, fees, and cage washings were collected.

The results of the study are given in Table 2E for Compound B; Table 2F for Zn-free Compound B; and Table 2G for modified Compound B.

TABLE 2E

Tissue Distribution of ¹⁴ C as a Percent of Dose in Rats Receiving Compound B Containing ¹⁴ C-NDGA, Dermally								
Tissue	Mean (± S.D.) ^a Percent of the Dose of ¹⁴ C Found at Hours							
	4	24	48	72	96			
Organs (%) Skinsite (%)	1.92 (±1.34) 16.3 (±9.2)	4.41 (±1.44) 11.9 (±6.1)	4.31 (±2.5) 8.92 (±4.10)	3.38 (±0.52) 6.55 (±1.76)	1.87 (±1.1) 5.11 (±3.0			

' aN = 3

TABLE 2F

Tissue Di	stribution of 14 C as a	Percent of Dose in F		ree Compound B	Containing		
Tissue Mean (± S.D.) ^a Percent of the Dose of ¹⁴ C Found at Hours							
	4	24	48	72	96		
Organs (%) Skinsite (%)	1.16 (±0.92) 2.21 (±1.44)	2.29 (±2.26) 4.45 (±4.36)	2.38 (±1.73) 6.07 (±2.90)	7.39 (±10.05) 18.6 (±6.3)	7.73 (±10.9) 12.0 (±4.99)		

aN = 3

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TABLE 2G

Tissue Distribution of ¹⁴ C as a Percent of Dose in Rats Receiving Modified Compound B Containing ¹⁴ C-NDGA, Dermally							
Tissue	Mean (± S.D.) ^a Percent of the Dose of ¹⁴ C Found at Hours						
	4	24	48	72	96		
Organs (%) Skinsite (%)	4.35 (±3.06) 14.4 (±7.1)		4.41 (±3.57) 17.1 (±4.7)	2.54 (±2.05) 16.3 (±0.6)	0.97 (±0.19) 17.7 (±6.2)		

⁸N = 3

The above results show that the addition of the water-soluble zinc-containing compound causes the organic molecule to more quickly absorb into the skin in larger quantities, and to be retained in the skin 30 longer than when the zinc-containing compound is not present.

EXAMPLE 3

Antineoplastic Enhancement

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For the organic compounds listed, in the case of those having two or more benzene rings, 0.145 moles per 100 cc²s was used, and for hose having one or no benzene rings, 0.29 moles per 100 cc²s was used. The organic compound was measured into a clean vial and PEGO 400 was added with mixing until dissolved.

For the test compounds containing zinc chloride, zinc chloride was first dissolved in the PEGO 400 to prepara a stock solution containing 0.69% zinc chloride, and this solution was added with mixing to the vials containing the organic compounds being tested.

These compounds were tested with and without zinc chloride for their effectiveness as antitumor agents against senograffs of human breast adenocarcinoma, MA-I, grown in athyrinic (nude) mice of Balbić so background by intratumor injection according to the following protocol: each animal was inaculated intradermally on the dorsum near the nape of the neck with 0.05 ml of an MX-1 tumor homogenate. Tumor weights, in milligrams, were calculated from a measurement of the length, width and height in millimeters of the tumors using the formula (L x W x H)/2. The animals were randomized in groups to ensure representation of smaller and larger tumors. The tumors were treated by intratumor injection with 0.05 ml of see also test composition. Each composition was tested utilizing five animals. The animals were treated only once. Results are set (toth in Table 3.

TABLE 3

5	Organic Compound Compound Known Anti-Cancer Agent	Tumor Free 60 Days	Prematur Death	a Tumor	Tumor Re-
	VP-16 (no Zn)	2	2	3	0
	VP-16 + 2n	5	0	0	0
10	*1/5 VP-16 + 2n	3	0	2	2
	VM-26 (no Zn)	0	0	5	0
	VM-26 + Zn	5	0	0	0
15	*1/5 VM-26 + Zn	5	0	0	0
	4'-demethylepipo- dophyllotoxin (no 2	n) 1	0	4	3
20	4'-demethylepipo- dophyllotoxin + 2n	5	0	0	0
	diethylstilbestrol (no 2n)	0	2	5	0
25	diethylstil- bestrol + Zn	3	0	2	, 1
	dithranol (no Zn)	1	0	4	0
	dithranol + Zn	4	0	1	1
30	cyclophosphamide (no 2n)	0	0	5	0
	cyclophosphamide + Zn	3	2	0	0
35	mitomycin (no Zn)	1	4	2	0
	mitomycin + Zn	5	0	0	0
	daunomycin (no Zn)	3	2	5	3
	daunomycin + Zn	5	0	0	0
40	platinum cis-diamine dichloride (no Zn)	1	0	4	0
	platinum cis-diamine dichloride + Zn	- 5	0	0	0

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TABLE 3 (Cont.)

5	Organic Compound Compound Known Anti-Cancer Agent	Tumor Free 60 Days	Premature Death a	Tumor it Death	Tumor Re-
	adriamycin (no Zn)	0	1	2	•
10	*1/10 adriamycin (no Zn)	2	1	2	2
	adriamycin + 2n	4	0	1	1
	*1/10 adriamycin + Zn	1	3	1	1
15	allopurinol (no Zn)	** 0		5	
	*5/2 allopurinol (no Zn)	0	0	5	0
20	allopurinol + Zn	1	0	4	4
	*5/2 allopurinol + Zn	1	0	4	4

- * Dosage level decreased or increased as indicated.
- ** All sacrificed on day 25.

These results show the reduced toxicity and enhanced antineoplastic effectiveness achieved by the $_{30}$ addition of zinc ions.

EXAMPLE 4

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Antineoplastic Potentiation

Compositions and organic compounds not previously known as antineoplastic agents were prepared with and without zinc chloride according to the procedure of Example 3 and tested for their ability to eradicate tumors following the protocol described in Example 3. Results are set forth in Table 4.

TABLE 4

Organic Compound	Tumor Free 60 Days	Premature Death	Tumor at Death	Tumor Recurrence
3-tertbutylphenol (no Zn)	1	3	1	0
3-tertbutylphenol + Zn	5	0	0	0
4-tertbutylphenol (no Zn)	5	0	0	0
4-tertbutylphenol + Zn	5	0	0	0
p-hydroxycinnamic acid (no Zn)	1	0	4	0
p-hydroxycinnamic acid + Zn	4	1	0	0
norisogualacin (no Zn)	2	1	1	0
norisogualacin + Zn	5	0	0	0
dl-NDGA (no Zn)	4	0	1	0
di-NDGA + Zn	5	0	0	0
azelaic acid (no Zn)	1	0	4	0
azelaic acid + Zn	5	0	0	0
1-(3,4-diacetoxyphenyl)-4-phenylbuta-1,3-diene (no Zn)	1	0	4	0
1-(3,4-diacetoxyphenyl)-4-phenylbuta-1,3-diene + Zn	3	0	2	2
1,4-bis(3,4-dihydroxyphenethyl)benzene (no Zn)	2	0	3	3
1,4-bis(3,4-dihydroxyphenethyl)benzene + Zn	2	0	3	3

These results show that the antineoplastic activity of drugs can be increased to worthwhile levels while at the same time reducing toxicity utilizing zinc ions.

EXAMPLES 5-10

Enhancement of Topical Antineoplastic Agents

In Examples 5-10, wherein reference is made to the testing of mixtures for antitumor activity against B if melanoms and Sacroma-180 solid tumor growth in mice, the following procedures were utilized. To the oxfort that a particular example modified the procedure, such modification will be indicated in the particular example.

Both types of tumors were grown intrademally or subcutaneously in the mice. The B-16 melanoma was grown in BOR mice. Each mouse was injected intradermally with about 0.01 ml of a saline suspension containing about 1 x 10° calls of the tumor cells per 0.01 ml into a preshaven area on the back of the neck of the mouse. The tumors were allowed to grow until they had an approximate size of about 25-100 mg, calculated by the length of the tumor multiplied by the width and height of the tumor seasured in millimeters and dividing the product by two. On the first day of teatment, the animals with tumor sizes outside of the size range were culled and the remaining animals were randomy divided into control and test groups. When the tumors had reached the appropriate size, usually at about day six, the tumors were punctured uniformly and then treated with either a test compound or a control by topical application to the surface of the tumor. Generally, two topical applications were made 24 hours apart. The materials were applied to obtain from about a 1 to about 2 mm containg over the surface of the tumor. The animals were thereafter observed and their weights and the size of their tumors were periodically measured.

The results of each of the experiments include the following:

- (a) the starting number starting number (n) of animals within a treatment group of an experiment;
- (b) the average tumor size in milligrams of the animals treated with the mixture and the average tumor size of the control animals:
 - (c) the ratio multiplied by 100 of the average size of the tumors of the treated animals to that of the control animals (T/C), wherein T = average size of treated mice and C = average tumor size of control mice:
- (d) the percentage of both created and control animals clear of tumor; and
- (e) the percentage of animals of the original number surviving.

The later three measurements for a particular experiment were all taken at the same time and range generally from 21 to about 33 days after tumor innoculation. A T/C value of 42 or less is indicative of activity, In all of the following tables for Examples 5-10, the control results are given in parenthesis ().

EXAMPLE 5

Two formulations of meso-NDGA were prepared in a PEGO base. Their compositions are as set forth in Table 5.

TABLE 5

Mixture	NDGA (meso)	H ₂ O	EtOH	PEGO 3350
1	3.6	24.3	48.5	23.4
2	6.9	0	0	93.1

Mixture No. 1 was prepared by dissolving the NDGA in absolute ethanol by warming and stirring; thereafter, the water was added slowly to the NDGA solution. The mixture was heated to evaporate sufficient solvent to obtain a mixture of about 130% of the weight of the NDGA, and was then incorporated into the PEGO base. Mixture No. 2 was made by simply dissolving the NDGA in the PEGO base with warming and stirring.

EXAMPLE 6

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The NDGA formulations of Example 5 were tested for potential antitumor activity against B-16 mananoma grown in mice. The procedure utilized was that previously described. The results are given below in Table 6.

TABLE 6

n	T/C1	Tumor Size (Control)	% Clear (Control)	% Survival (Control)	
9	87 114	954±698 (1091±547) 748±621 (645±335)	0 (0) 0 (0)	33 (80) 77 (77)	
10*	24	138±99 575±270	20 (0)	100 (60)	

¹T/C ratio was calculated between days 21-24.

EXAMPLE 7

Several formulations of zinc chloride in a PEGO base were prepared by first dissolving the zinc chloride in water and then mixing the zinc chloride solution into the PEGO base. The formulations had approximately the following weight/weight percent compositions as set forth in Table 7.

TABLE 7

Mixture	ZnCl ₂	H ₂ O	PEGO
3	41.9	11.6	46.5
4	30	16	53.8
5	14.6	4.1	81.3
6	28.6	7.9	63.5
7	46.1	13.8	40
8	15	8	77
9	13.8	9.2	77
10	5.5	3.7	90.8

[&]quot;Animal treated once via a 0.05 ml. intratumor injection on day 6 after tumor inoculation

EXAMPLE 8

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Mixtures of Example 7 were tested for potential antitumor activity against B-16 melanoma and S-180 solid tumor grown in mice in accordance with the procedures previously described. The results are given in Table 8.

TABLE 8

	Mixture	n	T/C ¹	Tumor Size (Control)	Clear (Control)	Survival (Control)
15	3	8	51	559±476 (1095±360)	25 (10)	100 (100)
20	3	10	40	366±345 (934±656)	11 (10)	90 (100)
	6	8	48	524±462 (1095±360)	12 (10)	100 (100)
25	6	10	115	1074±687 (934±656)	10 (10)	100 (100)
	7	5	0	0 (752±511)	100	80 (100)
30	7	10	0	0 (550±184)	100	90 (100)
35	7	10	0	0 (997±421)	100	100
	10	8	74	815±472 (1095±360)	0 (10)	87 (100)

¹T/C ratio calculated between days 20-23.

TABLE 8 (Cont.)

5 * Clear % Survival Tumor Size T/C1 (Control) (Control) (Control) Mixture n 60 100 6 162+253 3 10 10 (100) (2505±1844) (0) 100 106±114 30 4 10 16 (644±342) (0) (70)15 47 1191±764 22 5 10 (100) (0) (2505±1844) 100 94±217 80 6 10 4 (0) (100)(2502±1844) 20 37 88 169±216 7 9 15 (1091±547) (0) (80) 88 ٥ 100 0 25 7 9 . (77) (654±335) (0) 90 100 31±100 2 7 10 (80) (1805±968 (0) 30 40 43 277+209 10 8 10 (70) (644±342) (0) 80 90 25±63 10 1.0 9 (70) (80) (1733±2254) 35

1 T/C ratio calculated between days 21 and 25.

EXAMPLE 9

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Mixtures of zinc chloride, EDTA and NDGA were prepared and formulated in a PEGO base. The mixtures were prepared by dissolving the NDGA and EDTA in a portion of the PEGO base by warning and 4s sturing until dissolved. The zinc chloride was dissolved in water and warmed. The warm zinc chloride solution was added to the warm PEGO containing the NDGA and EDTA and stirred until cooled to room temperature. The composition of the mixtures is given in approximate weight/weight percentage as set forth in Table 9.

TABLE 9

Mixture	ZnCl₂	NDGA	EDTA	H ₂ O	PEGO	
11	27.5	6.9	14.7	18.3	32.6	
12	28	6.81	14.7	18.2	32.9	l

1 - T/C ratio calculated at day 24 except for Mixture 11 which was calculated at day 21.

EXAMPLE 10

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The mixtures of Example 9 were tested for their potential antitumor activities against B-16 melanomas proving in mice in accordance with the procedure previously described. The results are given in Table 10.

TABLE 10

B-16						
Mixture	n	T/C1	Tumor Size (Control)	% Clear (Control)	% Survival (Control)	
11	10	8	51±118 (711±286)	70 (0)	100 (100)	
12	10	0	0 (711±286)	60 (0)	100 (100)	

1 - T/C ratio calculated at day 24 except for Mixture 11 which was calculated at day 21.

The foregoing Examples 5-10 show the potentiating effect of zinc chloride on antineoplastic agents topically applied, showing improvement over the antineoplastic activity of zinc chloride alone, and comparable amounts of NDGA even when the NDGA is injected into the tumor.

EXAMPLE 11

Appropriate human and animal models are chosen for the disease conditions treated by the following compounds as herinabove described, and the compounds tested to determine effective and toxic dosages with and without the addition of equimolar amounts of zinc chloride, zinc iodide, zinc bromide, zinc sulfate, zinc nitrate, zinc stearate, and zinc acetate. From the results the Therapeutic Index, equivalent to ratio of toxic to effective dosage is calculated, and this Index for the compounds with and without zinc additives compared to show an increase in Therapeutic Index for compounds containing zinc additives as compared to compounds without such additives: NDGA (nordihydroguaiaretic acid), VP-16 (epipodophyllotoxin beta-D ethylidene glucopyranoside--etoposide), VM-26 (epipodophyllotoxin beta-D thenylidene glucopyranoside--teniposide), 4'demethyl epipodophyllotoxin, diethylstilbestrol, dithranol, cyclophosphamide, mitomycin, daunomycin, platinum cis-diamine-dichloride, adriamycin, allopurinol, 5-fluorcuracil, methotrexate, dexamethasone, hydrocortisone, hydrocortisone acetate, hydroxy hydrocortisone, hydrocortisone valerate, triamcinolone acetonide, triamcinolone hexacetonide, amcinonide, fluocinolone acetonide, fluocinolone flurandrenolide, difluorasone diacetate, betamethasone dipropionate, betamethasone, betamethasone benzoate, betamethasone valerate, halcinonide, desoximethasone, desonide, prednisolone, clocortolone pivalate, haloprogin, iodochloro, miconazole nitrate, tolnaftate, thiabendazole, chloroxine, amphotericin, candicin, fungimycin, nystatin, chlordantoin, clotrimazole, ethonam nitrate, miconazole nitrate, pyrrolnitrin fezatione, ticlatone, tolnaftate, triacetin, dithiocarbamate, thiourea, thiocyantes; aromatic carboxylic acids and the amides thereof, benzoic acid, salicylic acid, salicylic acid amide and anilide; aromatic sulfides, polysulfides, and sulfoxides, 5,5-dichloro-2,2-dihydroxydiphenylsulfide; quaternary ammonia and phosphonium compounds decamethylene-bis-(4-thiopyridine-methyl-tosylate; 8-hydroxyguinoline sulfate, halogenated quinolines, 7-iodo-8-hydroxy-quinoline-5-sulfonic acid, 5-chloro-7-iodo-8-hydroxy-quinoline, 5-5.7-dichlore-8-hydroxy-quinaldine. 5,7-diiodo-8-hydroxyquinoline, chloro-8-hydroxy-quinoline. decamethylene-bis (4-amino-quinaldium chloride): (2-dimethylamino-6-(beta-diaminoethoxy)-benzothiazote dihydrochloride; 1-(o-chloro-alpha-alpha-diphenyl-benzyl)-imidazole, 1-[o,p-dichloro-beta-(o,p-dichlorobenzyloxy)-phenethylimidazole; 2-phenylbenzimidazole, 2-furfurylbenzimidazole; 3,5-dibenzyltetrahydro-1,3,5thiadizine-2-thione; 5-nitro-2-furfuryl-3-chloropropionate; quinones, tetrachloro-p-benzoquinone, 1,4-naph-

thoquinone, phenanthraquinone; sulfonamide sulfones; aromatic diamidines, 2-hydroxystilbamidine, diamidinodiphenylamine, dioctyl sodium sulfosuccinate, miconazole nitrate, potassium sorbate, calcium propionate and sodium propionate, sodium lauryl sulfate, penicillin, cephalosporin, penicillinase, lincomycins, vancomycin, tetracyline, chloramphenicol, streptomycin, gramicidin, neomycin, polymyxin beta sulfate, 5 tetracycline, benzethonium chloride, gentamicin sulfate, nitrofurazone, benzalkonium chloride, hexylresorcinol, chloroxylenol, cloflucarban, carbolic acid (phenol), triclocarban, triclosan, glacial acetic acid, ascorbic acid, calcium pantothenate lactic acid, salicylic acid, cantharidin, podophyllin, acyclovir, benzalkonium chloride, alcohol, allantoin, anhydrous glycerin, benzocaine, camphor, carbamide peroxide, lanolin, menthol, petrolatum phenol, idoxuridine, amantadine, methisazone, cytarabine interferon, chloroform, ether, bacillus 10 calmette-querin, levamisole, benzimidazole, tetramisole, levamisole, isoquinoline, lidane, pyrethrin, piperonyl butoxide, malathion, crotamiton, benzoyl peroxide, resorcinol monoacetate, sulfur, povidone-iodine, phenol, fluocinolone acetonide, para-aminobenzoic acid, sodium thiosulfate, meclocyline sulfosalicylate, sodium sulfacetamide, tetracycline hydrochloride, 6-2 carbon aliphatic dicarboxylic acids, sulfurated lime, coal tar. juniper tar, pine tar, allantoin, saponated cresol, menthol, mercury oleate, phenol, methotrexate 4-4'-15 diaminodiphenyl sulfone, benzocaine, lidocaine, lidocaine hydrochloride, dibucaine, dibucaine hydrochloride, procaine, tetracaine, tetracaine hydrochloride, tronothane, dyclonine, dyclonine hydrochloride, pramoxine hydrochloride, benzyl alcohol, diperodon, butamben picrate, cyclomethycaine sulfate, dimethisoquin hydrochloride, N,N-dimethyl aspartic acid; N-N-dimethyl glutamic acid, trolamine salicylate, methyl salicylate; antipyrine, salicylamide, camphor, eugenol, eucalyptol, thymol, allyl isothiocyanate (mustard oil), capsicum 20 preparations, histamine dihydrochloride, methyl nicotinate, turpentine oil, diphenhydramine hydrochloride, phenyltoloxamine dihydrogen citrate, pyrilamine maleate, tripelennamine hydrochloride, tetanus antigen, diphtheria antigen, streptococcus antigen, tuberculin, Candida antigen, Trichophyton antigen, Proteus antigen, vitamins B3, B5, B6, A, D, and E, elastin, pregnenolone, hydroquinone, monobenzone, dioxybenzone, oxybenzone, padimate O, padimate A, aminobenzoic acid, cinoxate, diethanolamine p-methoxycin-25 namate, ethyl 4-[bis(hydroxypropyl)] aminobenzoate, ethylhexyl salicylate, glyceryl aminobenzoate, homosalate, lawsone, dihydroxyacetone, red petrolatum, and sulisobenzone.

Claims

- 30 1. The use of a water-soluble zinc-containing compound in a topical pharmaceutical composition containing a pharmacologically active agent to enchance the skin or mucous membrane penetration and retention of the pharmacologically active agent, and to reduce the transdermal flux through the skin and mucous membrane.
- The use as claimed in claim 1 wherein the pharmacologically active agent and the zinc-containing compound are present in approximately equimolar amounts.
 - The use as claimed in claim 1 or claim 2 wherein the concentration of zinc-containing compound is less than 35% by weight.
 - 4. The use as claimed in any one of claims 1 to 3 wherein the zinc-containing compound is a zinc salt.
 - The use as claimed in claim 4 wherein the zinc salt is a zinc halide, zinc sulfate, zinc nitrate, zinc acetate or zinc stearate.
 - 6. The use as claimed in claim 4 wherein the zinc salt is zinc chloride.
 - 7. The use as claimed in any one of the preceding claims wherein the pharmacologically active agent contains hydroxy, oxo, sulfhydryl, amine, carboxyl, or other anionic groups, or combinations thereof, in conformations which allow complexation and/or chelation by zinc ions.
 - 8. The use as claimed in any one of the preceding claims wherein the pharmacologically active agent is an antineoplastic agent, immunopharmacological agent, steroid, artiviral agent, antinflammatory agent, antiturgal agent, antiparastic agent, pediculicide, acree treatment compound, antipsoriasis agent, loproey agent, anaesthetic agent, counteriritant, antihistamine, allergy diagnostic agent, vitamin, nutrient, cosmetic agent or sunscreen.
 - 9. The use as claimed in any one of claims 1 to 6 wherein the pharmacologically active agent is

triamcinolone, acetonide, clotrimazole, miconazole or a salt thereof, salicylic salt, 5-fluorouracii, nordillydrogualaretic acid, vitamin A, retinoid, acyclovir, dapsone, hydrocortisone, tetracycline, urea or methotrexate.

5 10. A topical non-ocular pharmaceutical composition for application to the skin or mucous membrane having enhanced skin and mucous membrane penetration and retention, which composition comprises a pharmacologically active agent and a water-soluble zinc-containing compound wherein the pharmacologically active agent is VP-16 epipodophlylotxin beta-D ethylidene glucopyranoside--etoposide); VM-26 epipodophyllotoxin beta-D thenylidene glucopyranoside-teniposide); 4*-demethyl epipodophyllotoxin; 10 diethylstilbestrol; dithranol; cyclophosphamide; mitomycin; daunomycin; platinum cis-diamine-dichloride; adriamycin; allopurinol; haloprogin; iodochloro; tolnaftate, thiabendazole, chloroxine, amphotericin, candicin, fungimycin, nystatin, chlordantoin, clotrimazole, ethonam nitrate, miconazole nitrate, pyrrolnitrin, fezatione, ticalatone, triacetin, dithiocarbamate, thiourea, thiocyantes, aromatic carboxylic acids and the amides thereof, benzoic acid, salicylic acid amine and anilide, aromatic sulfides, polysulfides, and sulfoxides, 5.5.-dichloro-2, 2-dihydroxydiphenylsulfide; quarternary ammonia and phosphonium compounds, decamethylenebis-(4-thio-pyridine-methyl-tosylate, 8-hydroxyguinoline sulfate, halogenated quinolines, 7-iodo-8-hydroxy-quinoline-5-sulfonic acid, 5-chloro-7-iodo-8-hydroxy-quinoline, 5-chloro-8hydroxy-quinoline, 5,7,-dichloro-8-hydroxyquinaldine, 5,7-diiodo-8-hydroxyquinoline, decamethylene-bis (4-amino-quinaldium chloride); benzothiazole derivates, (2-dimethylamino-6-(beta-diaminoethoxy)-benzothiazole dihydrochloride; imidazole derivatives, 1-(o-chloro-alpha-alphadiphenylbenzyl)-imidazole, 1-20 [o,p-dichloro-beta-(o,p-dichlorobenzyloxy)-phenethylimi -dazole], 2-phenylbenzimidazole, 2-furfurylben-3,5-dibenzyltetrahydro-1,3,5-thiadizine-2-thione; 5-nitro-2-furfuryl-3-chloropropionate; quinones, tetrachloro-p-benzoquinoine, 1,4-naphthoquinone, phenanthraquinone; sulfonamides and sulfones; aromatic diamidines, 2-hydroxystilbamidine, diamidinodiphenylamine; dioctyl sodium sulfosuc-25 cinate, potassium sorbate, proprionate compounds, sodium lauryl sulfate; sulfonomides, penicillins, cephalosporins, penicillinase, lincomycins, vacomycins, tetracyline, chloram-phenicols, streptomycins, gramicidin, neomycin, poly-myxin beta sulfate, benzethonium chloride, gentamicin sulfate, nitrofurazone, benzalkonium chloride, hexylresorcinol, chloroxylenol, cloflucarban, triclocarban, triclocarban, glacial acetic acid, calcium panto-thenate, lactic acid, cantharidin, podophyllin, acyclovir, benzalkonium chloride, alcohol, anhydrous glycerin, benzocaine, camphor, carbamide peroxide, lanolin, menthol, petrolatum, urea, idoxuridine, amatadine, methisazone, cytarabine, interferon, chloroform, ether, bacillus calmette-guerin, levamisole; benzoyl peroxide, resorcinol monoacetate, azelaic acid, adipic acid, aliphatic dicarboxylic acids, sulfur, povidone-iodine, salicyclic acid, fluocinolone acetonide, para-aminobenzoic acid, sodium thiosulfate, meclocyline sulfosalicylate, sodium sulfacetamide, sulfurated lime; saponated cresol, menthol, mercury oleate, anthralin, methotrexate; N,N-dimethyl aspartic acid; N-N-35 dimethyl glutamic acid, antipyrine, camphor, eugenol, eucalyptol, thymol, allyl isothiocyanate (mustard oil) capsicum preparations, histamine dihydrochloride, methyl nicotinate, turpentine oil, diphenydramine hydrochloride, phenyltoloxamine dihydrogen citrate, pyrilamine maleate, or tripelennamine hydrochloride.

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